

**PARMETIS\***  
**Parallel Graph Partitioning and Sparse Matrix Ordering**  
**Library**  
**Version 3.1**

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# 1 Introduction

**PARMETIS** is an MPI-based parallel library that implements a variety of algorithms for partitioning and repartitioning unstructured graphs and for computing fill-reducing orderings of sparse matrices. **PARMETIS** is particularly suited for parallel numerical simulations involving large unstructured meshes. In this type of computation, **PARMETIS** dramatically reduces the time spent in communication by computing mesh decompositions such that the numbers of interface elements are minimized.

The algorithms in **PARMETIS** are based on the multilevel partitioning and fill-reducing ordering algorithms that are implemented in the widely-used serial package **METIS** [5]. However, **PARMETIS** extends the functionality provided by **METIS** and includes routines that are especially suited for parallel computations and large-scale numerical simulations. In particular, **PARMETIS** provides the following functionality:

- Partition unstructured graphs and meshes.
- Repartition graphs that correspond to adaptively refined meshes.
- Partition graphs for multi-phase and multi-physics simulations.
- Improve the quality of existing partitionings.
- Compute fill-reducing orderings for sparse direct factorization.
- Construct the dual graphs of meshes

The rest of this manual is organized as follows. Section 2 briefly describes the differences between this version and the previous major release. Section 3 describes the various algorithms that are implemented in **PARMETIS**. Section 4 describes the format of the basic parameters that need to be supplied to the routines. Section 5 provides a detailed description of the calling sequences for the major routines in **PARMETIS**. Finally, Section 6 describes software and hardware requirements and provides contact information.

## 2 What is New in This Version

**PARMETIS**, Version 3.x contains a number of changes over the previous major release (Version 2.x). These changes include the following:

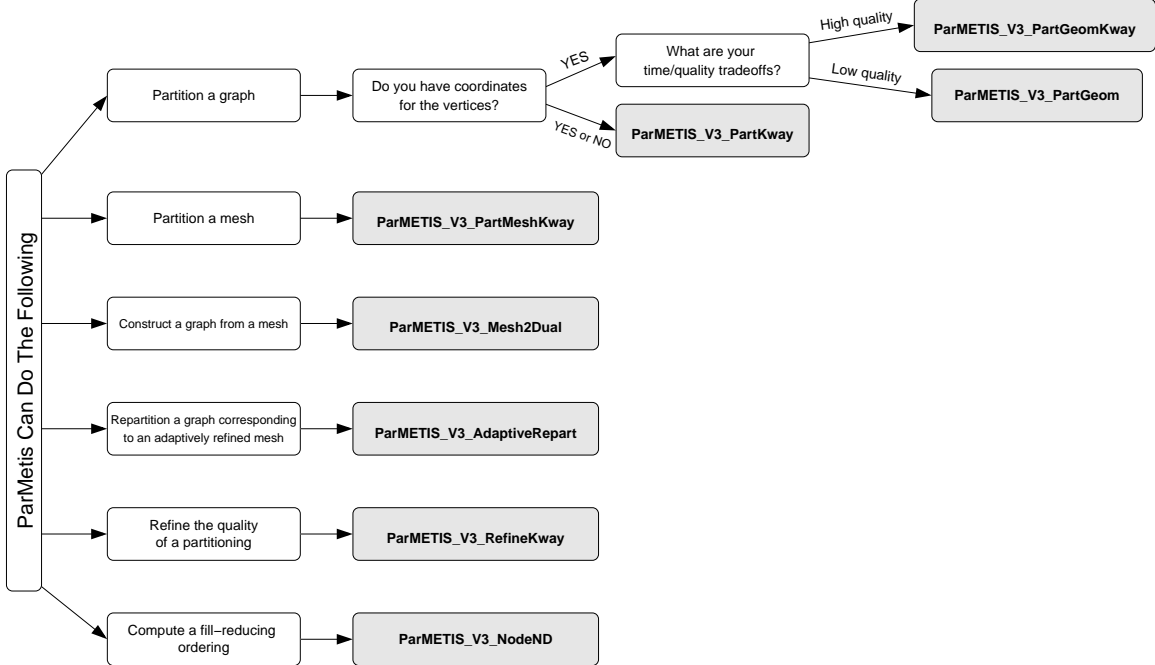
- The names and calling sequence of all the routines have changed due to expanded functionality that has been provided in this release. Table 1 shows how the names of the various routines map from version to version. Note that Version 3.0 is fully backwards compatible with all previous versions of **PARMETIS**. That is, the old API calls have been mapped to the new routines. However, the expanded functionality provided with this release is only available by using the new calling sequences.
- The four adaptive repartitioning routines: `ParMETIS_RepartLDiffusion`, `ParMETIS_RepartGDiffusion`, `ParMETIS_RepartRemap`, and `ParMETIS_RepartMLRemap` have been replaced by a (single) implementation of a unified repartitioning algorithm [15], `ParMETIS_V3_AdaptiveRepart`, that combines the best features of the previous routines.
- Multiple vertex weights/balance constraints are supported for most of the routines. This allows **PARMETIS** to be used to partition graphs for multi-phase and multi-physics simulations.
- In order to optimize partitionings for specific heterogeneous computing architectures, it is now possible to specify the target sub-domain weights for each of the sub-domains and for each balance constraint. This feature, for example, allows the user to compute a partitioning in which one of the sub-domains is twice the size of all of the others.
- The number of sub-domains has been de-coupled from the number of processors in both the static and the adaptive partitioning schemes. Hence, it is now possible to use the parallel partitioning and repartitioning algorithms to compute a  $k$ -way partitioning independent of the number of processors that are used. Note that Version 2.0 provided this functionality for the static partitioning schemes only.
- Routines are provided for both directly partitioning a finite element mesh, and for constructing the dual graph of a mesh in parallel. In version 3.1 these routines have been extended to support mixed element meshes.

Version 1.0	Version 2.0	Version 3.0
PARKMETIS	ParMETIS_PartKway	ParMETIS_V3_PartKway
PARGKMETIS	ParMETIS_PartGeomKway	ParMETIS_V3_PartGeomKway
PARGMETIS	ParMETIS_PartGeom	ParMETIS_V3_PartGeom
PARGRMETIS	Not available	Not available
PARRMETIS	ParMETIS_RefineKway	ParMETIS_V3_RefineKway
PARUAMETIS	ParMETIS_RepartLDiffusion	
PARDAMETIS	ParMETIS_RepartGDiffusion	
Not available	ParMETIS_RepartRemap	ParMETIS_V3_AdaptiveRepart
Not available	ParMETIS_RepartMLRemap	
PAROMETIS	ParMETIS_NodeND	ParMETIS_V3_NodeND
Not available	Not available	ParMETIS_V3_PartMeshKway
Not available	Not available	ParMETIS_V3_Mesh2Dual

**Table 1:** The relationships between the names of the routines in the different versions of **PARMETIS**.

### 3 Algorithms Used in PARMETIS

PARMETIS provides a variety of routines that can be used to compute different types of partitionings and repartitionings as well as fill-reducing orderings. Figure 1 provides an overview of the functionality provided by PARMETIS as well as a guide to its use.



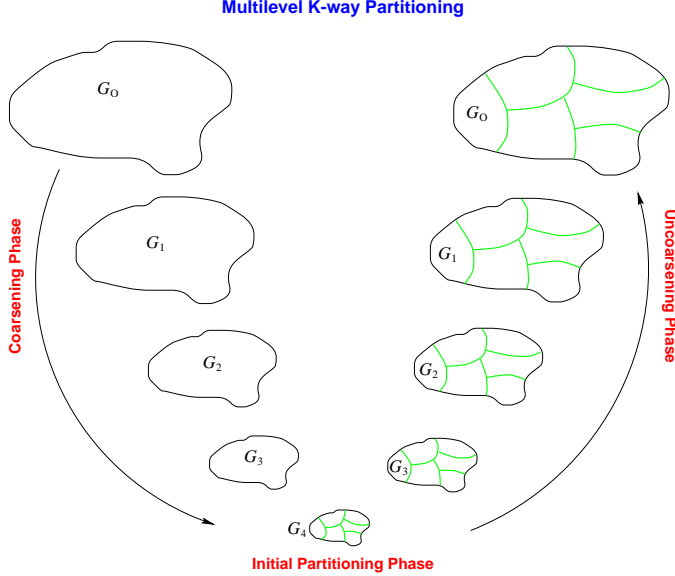
**Figure 1:** A brief overview of the functionality provided by PARMETIS. The shaded boxes correspond to the actual routines in PARMETIS that implement each particular operation.

#### 3.1 Unstructured Graph Partitioning

ParMETIS\_V3\_PartKway is the routine in PARMETIS that is used to partition unstructured graphs. This routine takes a graph and computes a  $k$ -way partitioning (where  $k$  is equal to the number of sub-domains desired) while attempting to minimize the number of edges that are cut by the partitioning (*i.e.*, the *edge-cut*). ParMETIS\_V3\_PartKway makes no assumptions on how the graph is initially distributed among the processors. It can effectively partition a graph that is randomly distributed as well as a graph that is well distributed<sup>1</sup>. If the graph is initially well distributed among the processors, ParMETIS\_V3\_PartKway will take less time to run. However, the quality of the computed partitionings does not depend on the initial distribution.

The parallel graph partitioning algorithm used in ParMETIS\_V3\_PartKway is based on the serial multilevel  $k$ -way partitioning algorithm described in [6, 7] and parallelized in [4, 14]. This algorithm has been shown to quickly produce partitionings that are of very high quality. It consists of three phases: graph coarsening, initial partitioning, and uncoarsening/refinement. In the graph coarsening phase, a series of graphs is constructed by collapsing together

<sup>1</sup>The reader should note the difference between the terms *graph distribution* and *graph partition*. A partitioning is a mapping of the vertices to the processors that results in a distribution. In other words, a partitioning specifies a distribution. In order to partition a graph in parallel, an initial distribution of the nodes and edges of the graph among the processors is required. For example, consider a graph that corresponds to the dual of a finite-element mesh. This graph could initially be partitioned simply by mapping groups of  $n/p$  consecutively numbered elements to each processor where  $n$  is the number of elements and  $p$  is the number of processors. Of course, this naive approach is not likely to result in a very good distribution because elements that belong to a number of different regions of the mesh may get mapped to the same processor. (That is, each processor may get a number of small sub-domains as opposed to a single contiguous sub-domain). Hence, you would want to compute a new high-quality partitioning for the graph and then redistribute the mesh accordingly. Note that it may also be the case that the initial graph is well distributed, as when meshes are adaptively refined and repartitioned.



**Figure 2:** The three phases of multilevel  $k$ -way graph partitioning. During the coarsening phase, the size of the graph is successively decreased. During the initial partitioning phase, a  $k$ -way partitioning is computed. During the multilevel refinement (or uncoarsening) phase, the partitioning is successively refined as it is projected to the larger graphs.  $G_0$  is the input graph, which is the finest graph.  $G_{i+1}$  is the next level coarser graph of  $G_i$ .  $G_4$  is the coarsest graph.

adjacent vertices of the input graph in order to form a related coarser graph. Computation of the initial partitioning is performed on the coarsest (and hence smallest) of these graphs, and so is very fast. Finally, partition refinement is performed on each level graph, from the coarsest to the finest (*i.e.*, original graph) using a KL/FM-type refinement algorithm [2, 9]. Figure 2 illustrates the multilevel graph partitioning paradigm.

Comparisons performed in [7] have shown that serial multilevel  $k$ -way partitioning is over 50 times faster than multilevel spectral bisection [12] while producing partitionings that cut 10% to 50% fewer edges. Our experiments on a 1024-processor Cray T3E have shown that **ParMETIS\_V3\_PartKway** can partition a 500-million element 3D mesh in well under a minute!

Recall that we mentioned that if the graph is well distributed among the processors **ParMETIS\_V3\_PartKway** runs faster. In fact, our experiments have shown that when we use **ParMETIS\_V3\_PartKway** to partition a graph that is distributed according to the partitioning produced by an earlier call to **ParMETIS\_V3\_PartKway**,<sup>2</sup> the amount of time required for this second partitioning is often reduced by a factor of two to four. The reason for this has to do with the inter-processor communication pattern of **ParMETIS\_V3\_PartKway**. If the graph is initially distributed randomly (*i.e.*, there are many interface vertices), each processor spends a lot of time communicating information about these interface vertices to many other processors. On the other hand, if the graph is well distributed, the number of interface vertices is much smaller (as is the number of other processors with whom each processor has to communicate), reducing the overall runtime of the partitioner. Of course, this is the chicken and egg problem. How can we initially distribute the graph nicely without having first partitioned it? We have developed one such method for partitioning graphs that correspond to finite element meshes. This is to quickly compute a fairly good initial partitioning using the coordinate values of the mesh. We can then redistribute the graph according to this initial partitioning and then call **ParMETIS\_V3\_PartKway** on the redistributed graph. **PARMETIS** provides the **ParMETIS\_V3\_PartGeomKway** routine for doing just this. Given a graph that is distributed among the processors and the coordinates of the vertices,<sup>3</sup> **ParMETIS\_V3\_PartGeomKway** quickly computes an initial partitioning using a space-filling curve method, redistributes the graph according to this partitioning, and then calls **ParMETIS\_V3\_PartKway** to compute the final high-quality partitioning. Our experiments have shown that **ParMETIS\_V3\_PartGeomKway** is often two times faster

<sup>2</sup>That is, we first call **ParMETIS\_V3\_PartKway** to find a good partitioning of a graph. Next we move the vertices of the graph according to the computed partitioning, and then call **ParMETIS\_V3\_PartKway** to partition this same, but newly distributed, graph.

<sup>3</sup>**ParMETIS\_V3\_PartGeomKway** requires the coordinates of the centers of each element.

than `ParMETIS_V3_PartKway`, and achieves identical partition quality.

**PARMETIS** also provides the `ParMETIS_PartGeom` function for partitioning unstructured graphs when coordinates for the vertices are available. `ParMETIS_PartGeom` computes a partitioning based only on the space-filling curve method. Therefore, it is extremely fast (often 5 to 10 times faster than `ParMETIS_PartGeomKway`), but it computes poor quality partitionings (it may cut 2 to 10 times more edges than `ParMETIS_PartGeomKway`). This routine can be useful for certain computations in which the use of space-filling curves is the appropriate partitioning technique (*e.g.*,  $n$ -body computations).

### 3.2 Partitioning Meshes Directly

**PARMETIS**, Version 3.0 also provides new routines that support the computation of partitionings and repartitionings given *meshes* (and not graphs) as inputs. In particular, `ParMETIS_V3_PartMeshKway` take a mesh as input and computes a partitioning of the mesh elements. Internally, `ParMETIS_V3_PartMeshKway` uses a mesh-to-graph routine and then calls the same core partitioning routine that is used by both `ParMETIS_V3_PartKway` and `ParMETIS_V3_PartGeomKway`.

**PARMETIS** provides no such routines for computing adaptive repartitionings directly from meshes. However, it does provide the routine `ParMETIS_V3_Mesh2Dual` for constructing a dual graph given a mesh, quickly and in parallel. Since the construction of the dual graph is in parallel, it can be used to construct the input graph for `ParMETIS_V3_AdaptiveRepart`.

Essentially, both `ParMETIS_V3_PartMeshKway` and `ParMETIS_V3_Mesh2Dual` take the burden of writing an efficient mesh-to-graph routine from the user. Our experiments have shown that this routine typically runs in about half the time that it takes for **PARMETIS** to compute a partitioning.

### 3.3 Partitioning Adaptively Refined Meshes

For large-scale scientific simulations, the computational requirements of techniques relying on globally refined meshes become very high, especially as the complexity and size of the problems increase. By locally refining and de-refining the mesh either to capture flow-field phenomena of interest [1] or to account for variations in errors [11], adaptive methods make standard computational methods more cost effective. The efficient execution of such adaptive scientific simulations on parallel computers requires a periodic repartitioning of the underlying computational mesh. These repartitionings should minimize both the inter-processor communications incurred in the iterative mesh-based computation and the data redistribution costs required to balance the load. Hence, adaptive repartitioning is a multi-objective optimization problem. **PARMETIS** provides the routine `ParMETIS_V3_AdaptiveRepart` for repartitioning such adaptively refined meshes. This routine assumes that the mesh is well distributed among the processors, but that (due to mesh refinement and de-refinement) this distribution is poorly load balanced.

Repartitioning algorithms fall into two general categories. The first category balances the computation by incrementally diffusing load from those sub-domains that have more work to adjacent sub-domains that have less work. These schemes are referred to as *diffusive schemes*. The second category balances the load by computing an entirely new partitioning, and then intelligently mapping the sub-domains of the new partitioning to the processors such that the redistribution cost is minimized. These schemes are generally referred to as *remapping schemes*. Remapping schemes typically lead to repartitionings that have smaller edge-cuts, while diffusive schemes lead to repartitionings that incur smaller redistribution costs. However, since these results can vary significantly among different types of applications, it can be difficult to select the best repartitioning scheme for the job.

Recently, we developed a Unified Repartitioning Algorithm [15] for adaptive repartitioning that combines the best characteristics of remapping and diffusion-based repartitioning schemes. A key parameter used by this algorithm is the *ITR Factor*. This parameter describes the ratio between the time required for performing the inter-processor communications incurred during parallel processing compared to the time to perform the data redistribution associated with balancing the load. As such, it allows us to compute a single metric that describes the quality of the repartitioning, even though adaptive repartitioning is a multi-objective optimization problem.

`ParMETIS_V3_AdaptiveRepart` is a parallel implementation of the Unified Repartitioning Algorithm. This is a

multilevel partitioning algorithm, and so, is in nature similar to the the algorithm implemented in `ParMETIS_V3_PartKway`. However, this routine uses a technique known as *local coarsening*. Here, only vertices that have been distributed onto the same processor are coarsened together. On the coarsest graph, an initial partitioning need not be computed, as one can either be derived from the initial graph distribution (in the case when sub-domains are coupled to processors), or else one needs to be supplied as an input to the routine (in the case when sub-domains are de-coupled from processors). However, this partitioning does need to be balanced. The balancing phase is performed on the coarsest graph twice by alternative methods. That is, optimized variants of remapping and diffusion algorithms [16] are both used to compute new partitionings. A quality metric for each of these partitionings is then computed (using the ITR Factor) and the partitioning with the highest quality is selected. This technique tends to give very good points from which to start multilevel refinement, regardless of the type of repartitioning problem or the value of the ITR Factor. Note that the fact that the algorithm computes two initial partitionings does not impact its scalability as long as the size of the coarsest graph is suitably small [8]. Finally, multilevel refinement is performed on the balanced partitioning in order to further improve its quality. Since `ParMETIS_V3_AdaptiveRepart` starts from a graph that is already well distributed, it is extremely fast. Experiments on a 1024-processor Cray T3E show that `ParMETIS_V3_AdaptiveRepart` is able to compute partitionings for a billion-element mesh in under a minute.

Appropriate values to pass for the ITR Factor parameter can easily be determined depending on the times required to perform (i) all inter-processor communications that have occurred since the last repartitioning, and (ii) the data redistribution associated with the last repartitioning/load balancing phase. Simply divide the first time by the second. The result is the correct ITR Factor. In case these times cannot be ascertained (*e.g.*, for the first repartitioning/load balancing phase), our experiments have shown that values between 100 and 1000 work well for a variety of situations.

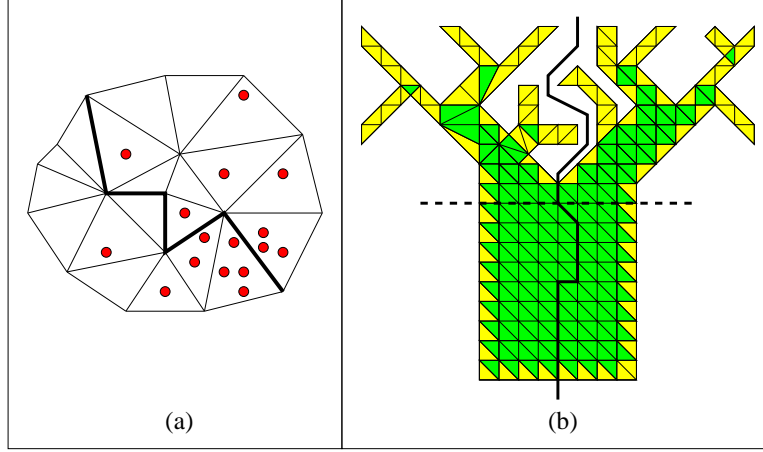
`ParMETIS_V3_AdaptiveRepart` can be used to load balance the mesh either before or after mesh adaptation. In the latter case, each processor first locally adapts its mesh, leading to different processors having different numbers of elements. `ParMETIS_V3_AdaptiveRepart` can then compute a partitioning in which the load is balanced. However, load balancing can also be done before adaptation if the degree of refinement for each element can be estimated *a priori*. That is, if we know ahead of time into how many new elements each old element will subdivide, we can use these estimations as the weights of the vertices for the graph that corresponds to the dual of the mesh. In this case, the mesh can be redistributed before adaption takes place. This technique can significantly reduce data redistribution times [10].

### 3.4 Partition Refinement

`ParMETIS_V3_RefineKway` is the routine provided by `PARMETIS` to improve the quality of an existing partitioning. Once a graph is partitioned and it has been redistributed accordingly, `ParMETIS_V3_RefineKway` can be called to compute a new partitioning that further improves the quality. Thus, like `ParMETIS_V3_AdaptiveRepart`, this routine assumes that the graph is already well distributed among the processors.

`ParMETIS_V3_RefineKway` can be used to improve the quality of partitionings that are produced by other partitioning algorithms (such as the technique discussed in Section 3.1 that is used in `ParMETIS_V3_PartGeom`). `ParMETIS_V3_RefineKway` can also be used repeatedly to further improve the quality of a partitioning. That is, we can call `ParMETIS_V3_RefineKway`, move the graph according to the partitioning, and then call `ParMETIS_V3_RefineKway` again. However, each successive call to `ParMETIS_V3_RefineKway` will tend to produce smaller improvements in quality.

Like `ParMETIS_V3_AdaptiveRepart`, `ParMETIS_V3_RefineKway` performs local coarsening. These two routines also use the same refinement algorithm. The difference is that `ParMETIS_V3_RefineKway` does not initially balance the partitioning on the coarsest graph, as `ParMETIS_V3_AdaptiveRepart` does. Instead, the assumption is that the graph is well distributed and the initial partitioning is balanced. Since `ParMETIS_V3_RefineKway` starts from a graph that is well distributed, it is very fast.



**Figure 3:** A computational mesh for a particle-in-cells simulation (a) and a computational mesh for a contact-impact simulation (b). The particle-in-cells mesh is partitioned so that both the number of mesh elements and the number of particles are balanced across the sub-domains. Two partitionings are shown for the contact-impact mesh. The dashed partitioning balances only the number of mesh elements. The solid partitioning balances both the number of mesh elements and the number of surface (lightly shaded) elements across the sub-domains.

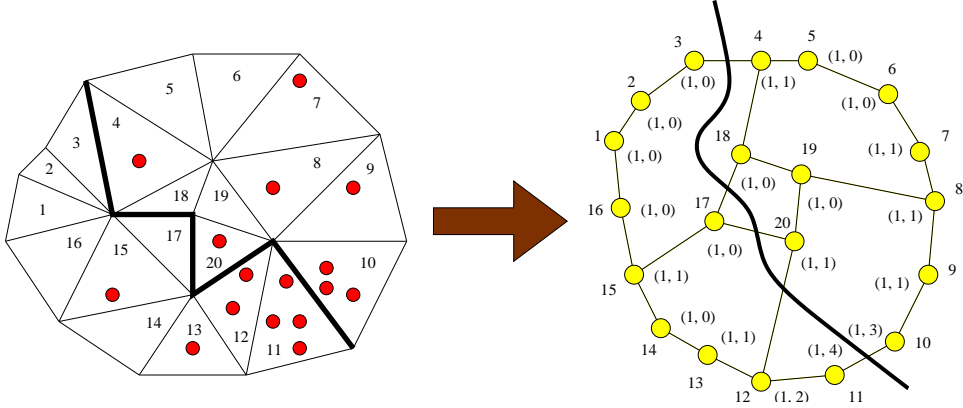
### 3.5 Partitioning for Multi-phase and Multi-physics Computations

The traditional graph partitioning problem formulation is limited in the types of applications that it can effectively model because it specifies that only a single quantity be load balanced. Many important types of multi-phase and multi-physics computations require that multiple quantities be load balanced simultaneously. This is because synchronization steps exist between the different phases of the computations, and so, each phase must be individually load balanced. That is, it is not sufficient to simply sum up the relative times required for each phase and to compute a partitioning based on this sum. Doing so may lead to some processors having too much work during one phase of the computation (and so, these may still be working after other processors are idle), and not enough work during another. Instead, it is critical that every processor have an equal amount of work from each phase of the computation.

Two examples are particle-in-cells [17] and contact-impact simulations [3]. Figure 3 illustrates the characteristics of partitionings that are needed for these simulations. Figure 3(a) shows a mesh for a particles-in-cells computation. Assuming that a synchronization separates the mesh-based computation from the particle computation, a partitioning is required that balances both the number of mesh elements and the number of particles across the sub-domains. Figure 3(b) shows a mesh for a contact-impact simulation. During the contact detection phase, computation is performed only on the surface (i.e., lightly shaded) elements, while during the impact phase, computation is performed on all of the elements. Therefore, in order to ensure that both phases are load balanced, a partitioning must balance both the total number of mesh elements and the number of surface elements across the sub-domains. The solid partitioning in Figure 3(b) does this. The dashed partitioning is similar to what a traditional graph partitioner might compute. This partitioning balances only the total number of mesh elements. The surface elements are imbalanced by over 50%.

A new formulation of the graph partitioning problem is presented in [6] that is able to model the problem of balancing multiple computational phases simultaneously, while also minimizing the inter-processor communications. In this formulation, a weight vector of size  $m$  is assigned to each vertex of the graph. The *multi-constraint graph partitioning problem* then is to compute a partitioning such that the edge-cut is minimized and that every sub-domain has approximately the same amount of each of the vertex weights. The routines `ParMETIS_V3_PartKway`, `ParMETIS_V3_PartGeomKway`, `ParMETIS_V3_RefineKway`, and `ParMETIS_V3_AdaptiveRepart` are all able to compute partitionings that satisfy multiple balance constraints.

Figure 4 gives the dual graph for the particles-in-cells mesh shown in Figure 3. Each vertex has two weights here. The first represents the work associated with the mesh-based computation for the corresponding element. (These are all ones because we assume in this case that all of the elements have the same amount of mesh-based work associated with them.) The second weight represents the work associated with the particle-based computation. This value is estimated



**Figure 4:** A dual graph with vertex weight vectors of size two is constructed from the particle-in-cells mesh from Figure 3. A multi-constraint partitioning has been computed for this graph, and this partitioning has been projected back to the mesh.

by the number of particles that fall within each element. A multi-constraint partitioning is shown that balances both of these weights.

A related graph partitioning problem formulation that is discussed in [13], allows edges to have multiple weights. We refer to this as the *multi-objective graph partitioning problem*. Multi-objective graph partitioning is applicable for tightly-coupled multi-physics computations involving multiple, spatially-overlapping meshes. Later versions of **ParMETIS** will support multiple edge weights.

### 3.6 Partitioning for Heterogeneous Computing Architectures

Complex, heterogeneous computing platforms, such as groups of tightly-coupled shared-memory nodes that are loosely connected via high bandwidth and high latency interconnection networks, and/or processing nodes that have complex memory hierarchies, are becoming more common, as they display competitive cost-to-performance ratios. The same is true of platforms that are geographically distributed. Most existing parallel simulation codes can easily be ported to a wide range of parallel architectures as they employ a standard messaging layer such as MPI. However, complex and heterogeneous architectures present new challenges to the scalable execution of such codes, since many of the basic parallel algorithm design assumptions are no longer valid. One way of alleviating this problem is to develop a new class of architecture-aware graph partitioning algorithms that optimally decompose computations given the architecture of the parallel platform. Ideally, this approach will alleviate the need for major restructuring of scientific codes.

We have taken the first steps toward developing architecture-aware graph-partitioning algorithms. These are able to compute partitionings that allow computations to achieve the highest levels of performance regardless of the computing platform. Specifically, we have enabled **ParMETIS\_V3\_PartKway**, **ParMETIS\_V3\_PartGeomKway**, **ParMETIS\_V3\_PartMeshKway**, **ParMETIS\_V3\_RefineKway**, and **ParMETIS\_V3\_AdaptiveRepart** to compute efficient partitionings for networks of heterogeneous processors. To do so, these routines require an additional array (`tpwgts`) to be passed as a parameter. This array describes the fraction of the total vertex weight each sub-domain should contain. For example, if you have a network of four processors, the first three of which are of equal processing speed, and the fourth of which is twice as fast as the others, the user would pass an array containing the values (0.2, 0.2, 0.2, 0.4). Note that by allowing users to specify target sub-domain weights as such, heterogeneous processing power can be taken into account when computing a partitioning. However, this does not allow us to take heterogeneous network bandwidths and latencies into account. Optimizing partitionings for heterogeneous networks is still the focus of ongoing research.

### 3.7 Computing Fill-Reducing Orderings

**ParMETIS\_V3\_NodeND** is the routine provided by **PARMETIS** for computing fill-reducing orderings, suited for

Cholesky-based direct factorization algorithms. **ParMETIS\_V3\_NodeND** makes no assumptions on how the graph is initially distributed among the processors. It can effectively compute fill-reducing orderings for graphs that are randomly distributed as well as graphs that are well distributed.

The algorithm implemented by **ParMETIS\_V3\_NodeND** is based on a multilevel nested dissection algorithm. This algorithm has been shown to produce low fill orderings for a wide variety of matrices. Furthermore, it leads to balanced elimination trees that are essential for parallel direct factorization. **ParMETIS\_V3\_NodeND** uses a multilevel node-based refinement algorithm that is particularly suited for directly refining the size of the separators. To achieve high performance, **ParMETIS\_V3\_NodeND** first uses **ParMETIS\_V3\_PartKway** to compute a high-quality partitioning and redistributes the graph accordingly. Next it proceeds to compute the  $\log p$  levels of the elimination tree concurrently. When the graph has been separated into  $p$  parts (where  $p$  is the number of processors), the graph is redistributed among the processor so that each processor receives a single subgraph, and a multiple minimum degree algorithm is used to order these smaller subgraphs.

## 4 Input and Output Formats used by **PARMETIS**

### 4.1 Format of the Input Graph

All of the graph routines in **PARMETIS** take as input the adjacency structure of the graph, the weights of the vertices and edges (if any), and an array describing how the graph is distributed among the processors. Note that depending on the application this graph can represent different things. For example, when **PARMETIS** is used to compute fill-reducing orderings, the graph corresponds to the non-zero structure of the matrix (excluding the diagonal entries). In the case of finite element computations, the vertices of the graph can correspond to nodes (points) in the mesh while edges represent the connections between these nodes. Alternatively, the graph can correspond to the dual of the finite element mesh. In this case, each vertex corresponds to an element and two vertices are connected via an edge if the corresponding elements share an edge (in 2D) or a face (in 3D). Also, the graph can be similar to the dual, but be more or less connected. That is, instead of limiting edges to those elements that share a face, edges can connect any two elements that share even a single node. However the graph is constructed, it is usually undirected.<sup>4</sup> That is, for every pair of connected vertices  $v$  and  $u$ , it contains both edges  $(v, u)$  and  $(u, v)$ .

In **PARMETIS**, the structure of the graph is represented by the compressed storage format (CSR), extended for the context of parallel distributed-memory computing. We will first describe the CSR format for serial graphs and then describe how it has been extended for storing graphs that are distributed among processors.

**Serial CSR Format** The CSR format is a widely-used scheme for storing sparse graphs. Here, the adjacency structure of a graph is represented by two arrays, `xadj` and `adjncy`. Weights on the vertices and edges (if any) are represented by using two additional arrays, `vwgt` and `adjwgt`. For example, consider a graph with  $n$  vertices and  $m$  edges. In the CSR format, this graph can be described using arrays of the following sizes:

`xadj`[ $n + 1$ ], `vwgt`[ $n$ ], `adjncy`[ $2m$ ], and `adjwgt`[ $2m$ ]

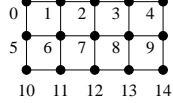
Note that the reason both `adjncy` and `adjwgt` are of size  $2m$  is because every edge is listed twice (*i.e.*, as  $(v, u)$  and  $(u, v)$ ). Also note that in the case in which the graph is unweighted (*i.e.*, all vertices and/or edges have the same weight), then either or both of the arrays `vwgt` and `adjwgt` can be set to `NULL`. `ParMETIS_V3_AdaptiveRepart` additionally requires a `vsize` array. This array is similar to the `vwgt` array, except that instead of describing the amount of work that is associated with each vertex, it describes the amount of memory that is associated with each vertex.

The adjacency structure of the graph is stored as follows. Assuming that vertex numbering starts from 0 (C style), the adjacency list of vertex  $i$  is stored in array `adjncy` starting at index `xadj`[ $i$ ] and ending at (but not including) index `xadj`[ $i + 1$ ] (in other words, `adjncy`[`xadj`[ $i$ ]] up through and including `adjncy`[`xadj`[ $i + 1$ ]-1]). Hence, the adjacency lists for each vertex are stored consecutively in the array `adjncy`. The array `xadj` is used to point to where the list for each specific vertex begins and ends. Figure 5(b) illustrates the CSR format for the 15-vertex graph shown in Figure 5(a). If the graph has weights on the vertices, then `vwgt`[ $i$ ] is used to store the weight of vertex  $i$ . Similarly, if the graph has weights on the edges, then the weight of edge `adjncy`[ $j$ ] is stored in `adjwgt`[ $j$ ]. This is the same format that is used by the (serial) **METIS** library routines.

**Distributed CSR Format** **PARMETIS** uses an extension of the CSR format that allows the vertices of the graph and their adjacency lists to be distributed among the processors. In particular, **PARMETIS** assumes that each processor  $P_i$  stores  $n_i$  consecutive vertices of the graph and the corresponding  $m_i$  edges, so that  $n = \sum_i n_i$ , and  $2 * m = \sum_i m_i$ . Here, each processor stores its local part of the graph in the four arrays `xadj`[ $n_i + 1$ ], `vwgt`[ $n_i$ ], `adjncy`[ $m_i$ ], and `adjwgt`[ $m_i$ ], using the CSR storage scheme. Again, if the graph is unweighted, the arrays `vwgt` and `adjwgt` can be set to `NULL`. The straightforward way to distribute the graph for **PARMETIS** is to take  $n/p$  consecutive adjacency lists from `adjncy` and store them on consecutive processors (where  $p$  is the number of processors). In addition, each

---

<sup>4</sup>Multi-constraint and multi-objective graph partitioning formulations [6, 13] can get around this requirement for some applications. These also allow the computation of partitionings for bipartite graphs, as well as for graphs corresponding to non-square and non-symmetric matrices.



(a) A sample graph

Description of the graph on a serial computer (serial MeTiS)

xadj	0 2 5 8 11 13 16 20 24 28 31 33 36 39 42 44
adjncy	1 5 0 2 6 1 3 7 2 4 8 3 9 0 6 10 1 5 7 11 2 6 8 12 3 7 9 13 4 8 14 5 11 6 10 12 7 11 13 8 12 14 9 13

(b) Serial CSR format

Description of the graph on a parallel computer with 3 processors (ParMeTiS)

Processor 0:	xadj	0 2 5 8 11 13
	adjncy	1 5 0 2 6 1 3 7 2 4 8 3 9
	vtxdist	0 5 10 15
Processor 1:	xadj	0 3 7 11 15 18
	adjncy	0 6 10 1 5 7 11 2 6 8 12 3 7 9 13 4 8 14
	vtxdist	0 5 10 15
Processor 2:	xadj	0 2 5 8 11 13
	adjncy	5 11 6 10 12 7 11 13 8 12 14 9 13
	vtxdist	0 5 10 15

(c) Distributed CSR format

**Figure 5:** An example of the parameters passed to **PARMETIS** in a three processor case. The arrays `vwgt` and `adjwgt` are assumed to be NULL.

processor needs its local `xadj` array to point to where each of its local vertices' adjacency lists begin and end. Thus, if we take all the local `adjncy` arrays and concatenate them, we will get exactly the same `adjncy` array that is used in the serial CSR. However, concatenating the local `xadj` arrays will not give us the serial `xadj` array. This is because the entries in each local `xadj` must point to their local `adjncy` array, and so, `xadj[0]` is zero for all processors. In addition to these four arrays, each processor also requires the array `vtxdist[p + 1]` that indicates the range of vertices that are local to each processor. In particular, processor  $P_i$  stores the vertices from `vtxdist[i]` up to (but not including) vertex `vtxdist[i + 1]`.

Figure 5(c) illustrates the distributed CSR format by an example on a three-processor system. The 15-vertex graph in Figure 5(a) is distributed among the processors so that each processor gets 5 vertices and their corresponding adjacency lists. That is, Processor Zero gets vertices 0 through 4, Processor One gets vertices 5 through 9, and Processor Two gets vertices 10 through 14. This figure shows the `xadj`, `adjncy`, and `vtxdist` arrays for each processor. Note that the `vtxdist` array will always be identical for every processor.

All five arrays that describe the distributed CSR format are defined in **PARMETIS** to be of type `idxtype`. By default `idxtype` is set to be equivalent to type `int` (*i.e.*, integers). However, `idxtype` can be made to be equivalent to a `short int` for certain architectures that use 64-bit integers by default. (Note that doing so will cut the memory usage and communication time required approximately in half.) The conversion of `idxtype` from `int` to `short` can be done by modifying the file `parmetis.h`. (Instructions are included there.) The same `idxtype` is used for the arrays that store the computed partitioning and permutation vectors.

When multiple vertex weights are used for multi-constraint partitioning, the  $c$  vertex weights for each vertex are stored contiguously in the `vwgt` array. In this case, the `vwgt` array is of size  $nc$ , where  $n$  is the number of locally-stored vertices and  $c$  is the number of vertex weights (and also the number of balance constraints).

## 4.2 Format of Vertex Coordinates

As discussed in Section 3.1, **PARMETIS** provides routines that use the coordinate information of the vertices to quickly pre-distribute the graph, and so, speedup the execution of the parallel  $k$ -way partitioning. These coordinates are specified in an array called `xyz` of single precision floating point numbers (*i.e.*, `float`). If  $d$  is the number of dimensions of the mesh (*i.e.*,  $d = 2$  for 2D meshes or  $d = 3$  for 3D meshes), then each processor requires an array of size  $d * n_i$ , where  $n_i$  is the number of locally-stored vertices. (Note that the number of dimensions of the mesh,  $d$ , is required as a parameter to the routine.) In this array, the coordinates of vertex  $i$  are stored starting at location `xyz[i * d]` up to (but not including) location `xyz[i * d + d]`. For example, if  $d = 3$ , then the x, y, and z coordinates of vertex  $i$  are stored at `xyz[3 * i]`, `xyz[3 * i + 1]`, and `xyz[3 * i + 2]`, respectively.

## 4.3 Format of the Input Mesh

The routine `ParMETIS_V3_PartMeshKway` takes a distributed mesh and computes its partitioning, while `ParMETIS_V3_Mesh2Dual` takes a distributed mesh and constructs a distributed dual graph. Both of these routines require an `e1mdist` array that specifies the distribution of the mesh elements, but that is otherwise identical to the `vtxdist` array. They also require a pair of arrays called `eptr` and `eind`, as well as the integer parameter `ncommonnodes`.

The `eptr` and `eind` arrays are similar in nature to the `xadj` and `adjncy` arrays used to specify the adjacency list of a graph but now for each element they specify the set of nodes that make up each element. Specifically, the set of nodes that belong to element  $i$  is stored in array `eind` starting at index `eptr[i]` and ending at (but not including) index `eptr[i + 1]` (in other words, `eind[eptr[i]]` up through and including `eind[eptr[i + 1] - 1]`). Hence, the node lists for each element are stored consecutively in the array `eind`. This format allows the specification of meshes that contain elements of mixed type.

The `ncommonnodes` parameter specifies the degree of connectivity that is desired between the vertices of the dual graph. Specifically, an edge is placed between two vertices if their corresponding mesh elements share at least  $g$  nodes, where  $g$  is the `ncommonnodes` parameter. Hence, this parameter can be set to result in a traditional dual graph (*e.g.*, a value of two for a triangle mesh or a value of four for a hexahedral mesh). However, it can also be set higher or lower for increased or decreased connectivity. `ParMETIS_V3_PartMeshKway` additionally requires an `e1mwgt` array that is analogous to the `vwgt` array.

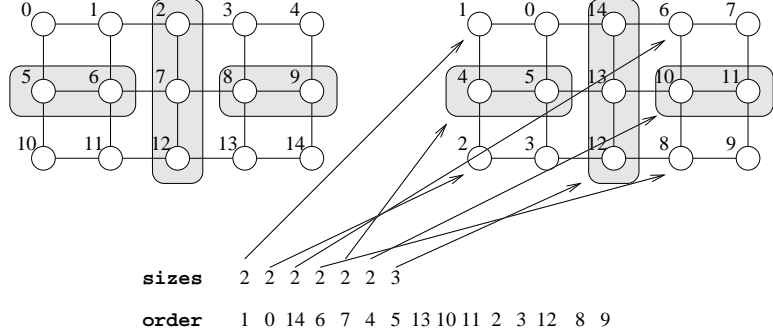
## 4.4 Format of the Computed Partitionings and Orderings

**Format of the Partitioning Array** The partitioning and repartitioning routines require that arrays (called `part`) of sizes  $n_i$  (where  $n_i$  is the number of local vertices) be passed as parameters to each processor. Upon completion of the **PARMETIS** routine, for each vertex  $j$ , the sub-domain number (*i.e.*, the processor label) to which this vertex belongs will have been written to `part[j]`. Note that **PARMETIS** does not redistribute the graph according to the new partitioning, it simply computes the partitioning and writes it to the `part` array.

Additionally, whenever the number of sub-domains does not equal the number of processors that are used to compute a repartitioning, `ParMETIS_V3_AdaptiveReport` requires that the previously computed partitioning be passed as a parameter via the `part` array. (This is also required whenever the user chooses to de-couple the sub-domains from the processors. See discussion in Section 5.2.) This is because the initial partitioning needs to be obtained from the values supplied in the `part` array. If the numbers of sub-domains and processors are equal, then the initial partitioning can be obtained from the initial graph distribution, and so this information need not be supplied. (In this case, for each processor  $i$ , every element of `part` would be set to  $i$ .)

**Format of the Permutation Array** Likewise, each processor running `ParMETIS_V3_NodeND` writes its portion of the computed fill-reducing ordering to an array called `order`. Similar to the `part` array, the size of `order` is equal to the number of vertices stored at each processor. Upon completion, for each vertex  $j$ , `order[j]` stores the new global number of this vertex in the fill-reducing permutation.

Besides the ordering vector, `ParMETIS_V3_NodeND` also returns information about the sizes of the different



**Figure 6:** An example of the ordering produced by ParMETIS\_V3\_NodeND. Consider the simple  $3 \times 5$  grid and assume that we have four processors. ParMETIS\_V3\_NodeND finds the three separators that are shaded. It first finds the big separator and then for each of the two sub-domains it finds the smaller. At the end of the ordering, the `order` vector concatenated over all the processors will be the one shown. Similarly, the `sizes` arrays will all be identical to the one shown, corresponding to the regions pointed to by the arrows.

sub-domains as well as the separators at different levels. This array is called `sizes` and is of size  $2p$  (where  $p$  is the number of processors). Every processor must supply this array and upon return, each of the `sizes` arrays are identical.

The format of this array is as follows. The first  $p$  entries of `sizes` starting from 0 to  $p - 1$  store the number of nodes in each one of the  $p$  sub-domains. The remaining  $p - 1$  entries of this array starting from `sizes`[ $p$ ] up to `sizes`[ $2p - 2$ ] store the sizes of the separators at the  $\log p$  levels of nested dissection. In particular, `sizes`[ $2p - 2$ ] stores the size of the top level separator, `sizes`[ $2p - 4$ ] and `sizes`[ $2p - 3$ ] store the sizes of the two separators at the second level (from left to right). Similarly, `sizes`[ $2p - 8$ ] through `sizes`[ $2p - 5$ ] store the sizes of the four separators of the third level (from left to right), and so on. This array can be used to quickly construct the separator tree (a form of an elimination tree) for direct factorization. Given this separator tree and the sizes of the sub-domains, the nodes in the ordering produced by ParMETIS\_V3\_NodeND are numbered in a postorder fashion. Figure 6 illustrates the `sizes` array and the postorder ordering.

## 4.5 Numbering and Memory Allocation

PARMETIS allows the user to specify a graph whose numbering starts either at 0 (C style) or at 1 (Fortran style). Of course, PARMETIS requires that same numbering scheme be used consistently for all the arrays passed to it, and it writes to the `part` and `order` arrays similarly.

PARMETIS allocates all the memory that it requires dynamically. This has the advantage that the user does not have to provide workspace. However, if there is not enough memory on the machine, the routines in PARMETIS will abort. Note that the routines in PARMETIS do not modify the arrays that store the graph (*e.g.*, `xadj` and `adjncy`). They only modify the `part` and `order` arrays.

## 5 Calling Sequence of the Routines in **PARMETIS**

The calling sequences of the **PARMETIS** routines are described in this section.

## 5.1 Graph Partitioning

**ParMETIS\_V3\_PartKway** (idxtype \*vtxdist, idxtype \*xadj, idxtype \*adjncy, idxtype \*vwgt, idxtype \*adjwgt,  
int \*wgtflag, int \*numflag, int \*ncon, int \*nparts, float \*tpwgts, float \*ubvec,  
int \*options, int \*edgecut, idxtype \*part, MPI\_Comm \*comm)

### Description

This routine is used to compute a  $k$ -way partitioning of a graph on  $p$  processors using the multilevel  $k$ -way multi-constraint partitioning algorithm.

### Parameters

**vtxdist** This array describes how the vertices of the graph are distributed among the processors. (See discussion in Section 4.1). Its contents are identical for every processor.

**xadj, adjncy**

These store the (local) adjacency structure of the graph at each processor. (See discussion in Section 4.1).

**vwgt, adjwgt**

These store the weights of the vertices and edges. (See discussion in Section 4.1).

**wgtflag** This is used to indicate if the graph is weighted. *wgtflag* can take one of four values:

- 0 No weights (*vwgt* and *adjwgt* are both NULL).
- 1 Weights on the edges only (*vwgt* is NULL).
- 2 Weights on the vertices only (*adjwgt* is NULL).
- 3 Weights on both the vertices and edges.

**numflag** This is used to indicate the numbering scheme that is used for the *vtxdist*, *xadj*, *adjncy*, and *part* arrays. *numflag* can take one of two values:

- 0 C-style numbering that starts from 0.
- 1 Fortran-style numbering that starts from 1.

**ncon** This is used to specify the number of weights that each vertex has. It is also the number of balance constraints that must be satisfied.

**nparts** This is used to specify the number of sub-domains that are desired. Note that the number of sub-domains is independent of the number of processors that call this routine.

**tpwgts** An array of size *ncon* x *nparts* that is used to specify the fraction of vertex weight that should be distributed to each sub-domain for each balance constraint. If all of the sub-domains are to be of the same size for every vertex weight, then each of the *ncon* x *nparts* elements should be set to a value of  $1/nparts$ . If *ncon* is greater than one, the target sub-domain weights for each sub-domain are stored contiguously (similar to the *vwgt* array). Note that the sum of all of the *tpwgts* for a give vertex weight should be one.

**ubvec** An array of size *ncon* that is used to specify the imbalance tolerance for each vertex weight, with 1 being perfect balance and *nparts* being perfect imbalance. A value of 1.05 for each of the *ncon* weights is recommended.

**options** This is an array of integers that is used to pass additional parameters for the routine. If *options[0]*=0, then the default values are used. If *options[0]*=1, then the remaining two elements of *options* are interpreted as follows:

**options[1]** This specifies the level of information to be returned during the execution of the algorithm. Timing information can be obtained by setting this to 1. Additional options for this parameter can be obtained by looking at the file *defs.h* in the ParMETIS-Lib directory. The numerical values there should be added to obtain the correct value. The default value is 0.

- options[2] This is the random number seed for the routine. The default value is 15.
- edgecut** Upon successful completion, the number of edges that are cut by the partitioning is written to this parameter.
- part** This is an array of size equal to the number of locally-stored vertices. Upon successful completion the partition vector of the locally-stored vertices is written to this array. (See discussion in Section 4.4).
- comm** This is a pointer to the MPI communicator of the processes that call PARMETIS. For most programs this will point to MPI\_COMM\_WORLD.

```
ParMETIS_V3_PartGeomKway (idxtype *vtxdist, idxtype *xadj, idxtype *adjncy, idxtype *vwgt, idxtype *adjwgt,
int *wgtflag, int *numflag, int *ndims, float *xyz, int *ncon, int *nparts,
float *tpwgts, float *ubvec, int *options, int *edgecut, idxtype *part,
MPI_Comm *comm)
```

## Description

This routine is used to compute a  $k$ -way partitioning of a graph on  $p$  processors by combining the coordinate-based and multi-constraint  $k$ -way partitioning schemes.

## Parameters

- vtxdist** This array describes how the vertices of the graph are distributed among the processors. (See discussion in Section 4.1). Its contents are identical for every processor.
- xadj, adjncy** These store the (local) adjacency structure of the graph at each processor. (See discussion in Section 4.1).
- vwgt, adjwgt** These store the weights of the vertices and edges. (See discussion in Section 4.1).
- wgtflag** This is used to indicate if the graph is weighted. *wgtflag* can take one of four values:
  - 0 No weights (vwgt and adjwgt are both NULL).
  - 1 Weights on the edges only (vwgt is NULL).
  - 2 Weights on the vertices only (adjwgt is NULL).
  - 3 Weights on both the vertices and edges.
- numflag** This is used to indicate the numbering scheme that is used for the *vtxdist*, *xadj*, *adjncy*, and *part* arrays. *numflag* can take one of two values:
  - 0 C-style numbering that starts from 0.
  - 1 Fortran-style numbering that starts from 1.
- ndims** The number of dimensions of the space in which the graph is embedded.
- xyz** The array storing the coordinates of the vertices (described in Section 4.2).
- ncon** This is used to specify the number of weights that each vertex has. It is also the number of balance constraints that must be satisfied.
- nparts** This is used to specify the number of sub-domains that are desired. Note that the number of sub-domains is independent of the number of processors that call this routine.
- tpwgts** An array of size *ncon* x *nparts* that is used to specify the fraction of vertex weight that should be distributed to each sub-domain for each balance constraint. If all of the sub-domains are to be of the same size for every vertex weight, then each of the *ncon* x *nparts* elements should be set to a value of  $1/nparts$ . If *ncon* is greater than one, the target sub-domain weights for each sub-domain are stored contiguously (similar to the *vwgt* array). Note that the sum of all of the *tpwgts* for a give vertex weight should be one.
- ubvec** An array of size *ncon* that is used to specify the imbalance tolerance for each vertex weight, with 1 being perfect balance and *nparts* being perfect imbalance. A value of 1.05 for each of the *ncon* weights is recommended.
- options** This is an array of integers that is used to pass parameters to the routine. Their meanings are identical to those of **ParMETIS\_V3\_PartKway**.
- edgecut** Upon successful completion, the number of edges that are cut by the partitioning is written to this parameter.

- part** This is an array of size equal to the number of locally-stored vertices. Upon successful completion the partition vector of the locally-stored vertices is written to this array. (See discussion in Section 4.4).
- comm** This is a pointer to the MPI communicator of the processes that call **PARMETIS**. For most programs this will point to **MPI\_COMM\_WORLD**.

**Note**

The quality of the partitionings computed by **ParMETIS\_V3\_PartGeomKway** are comparable to those produced by **ParMETIS\_V3\_PartKway**. However, the run time of the routine may be up to twice as fast.

**ParMETIS\_V3\_PartGeom** (idxtype \*vtxdist, int \*ndims, float \*xyz, idxtype \*part, MPI\_Comm \*comm)

### Description

This routine is used to compute a  $p$ -way partitioning of a graph on  $p$  processors using a coordinate-based space-filling curves method.

### Parameters

- vtxdist** This array describes how the vertices of the graph are distributed among the processors. (See discussion in Section 4.1). Its contents are identical for every processor.
- ndims** The number of dimensions of the space in which the graph is embedded.
- xyz** The array storing the coordinates of the vertices (described in Section 4.2).
- part** This is an array of size equal to the number of locally stored vertices. Upon successful completion stores the partition vector of the locally stored graph (described in Section 4.4).
- comm** This is a pointer to the MPI communicator of the processes that call ParMETIS. For most programs this will point to MPI\_COMM\_WORLD.

### Note

The quality of the partitionings computed by ParMETIS\_V3\_PartGeom are significantly worse than those produced by ParMETIS\_V3\_PartKway and ParMETIS\_V3\_PartGeomKway.

```
ParMETIS_V3_PartMeshKway (idxtype *elmdist, idxtype *eptr, idxtype *eind, idxtype *elmwgt,
int *wgtflag, int *numflag, int *ncon, int *ncommonnodes, int *nparts,
float *tpwgts, float *ubvec, int *options, int *edgecut, idxtype *part,
MPI_Comm *comm)
```

## Description

This routine is used to compute a  $k$ -way partitioning of a *mesh* on  $p$  processors. The mesh can contain elements of different types.

## Parameters

- elmdist** This array describes how the elements of the mesh are distributed among the processors. It is analogous to the `vtxdist` array. Its contents are identical for every processor. (See discussion in Section 4.3).
- eptr, eind** These arrays specifies the elements that are stored locally at each processor. (See discussion in Section 4.3).
- elmwgt** This array stores the weights of the elements. (See discussion in Section 4.3).
- wgtflag** This is used to indicate if the graph is weighted. `wgtflag` can take one of four values:
  - 0 No weights (`vwgt` and `adjwgt` are both NULL).
  - 1 Weights on the edges only (`vwgt` is NULL).
  - 2 Weights on the vertices only (`adjwgt` is NULL).
  - 3 Weights on both the vertices and edges.
- numflag** This is used to indicate the numbering scheme that is used for the `elmdist`, `elements`, and `part` arrays. `numflag` can take one of two values:
  - 0 C-style numbering that starts from 0.
  - 1 Fortran-style numbering that starts from 1.
- ncon** This is used to specify the number of weights that each vertex has. It is also the number of balance constraints that must be satisfied.
- ncommonnodes** This parameter determines the degree of connectivity among the vertices in the dual graph. Specifically, an edge is placed between any two elements if and only if they share at least this many nodes. This value should be greater than zero, and for most meshes a value of two will create reasonable dual graphs. However, depending on the type of elements in the mesh, values greater than two may also be valid choices. For example, for meshes containing only triangular, tetrahedral, hexahedral, or rectangular elements, this parameter can be set to two, three, four, or two, respectively.  
Note that setting this parameter to a small value will increase the number of edges in the resulting dual graph and the corresponding partitioning time.
- nparts** This is used to specify the number of sub-domains that are desired. Note that the number of sub-domains is independent of the number of processors that call this routine.
- tpwgts** An array of size `ncon` x `nparts` that is used to specify the fraction of vertex weight that should be distributed to each sub-domain for each balance constraint. If all of the sub-domains are to be of the same size for every vertex weight, then each of the `ncon` x `nparts` elements should be set to a value of  $1/nparts$ . If `ncon` is greater than one, the target sub-domain weights for each sub-domain are stored contiguously (similar to the `vwgt` array). Note that the sum of all of the `tpwgts` for a give vertex weight should be one.

- ubvec** An array of size `ncon` that is used to specify the imbalance tolerance for each vertex weight, with 1 being perfect balance and `nparts` being perfect imbalance. A value of 1.05 for each of the `ncon` weights is recommended.
- options** This is an array of integers that is used to pass parameters to the routine. Their meanings are identical to those of `ParMETIS_V3_PartKway`.
- edgecut** Upon successful completion, the number of edges that are cut by the partitioning is written to this parameter.
- part** This is an array of size equal to the number of locally-stored vertices. Upon successful completion the partition vector of the locally-stored vertices is written to this array. (See discussion in Section 4.4).
- comm** This is a pointer to the MPI communicator of the processes that call `PARMETIS`. For most programs this will point to `MPI_COMM_WORLD`.

## 5.2 Graph Repartitioning

```
ParMETIS_V3_AdaptiveReport (idxtype *vtxdist, idxtype *xadj, idxtype *adjncy, idxtype *vwgt, idxtype *vsize,
                           idxtype *adjwgt, int *wgtflag, int *numflag, int *ncon, int *nparts, float *tpwgts,
                           float *ubvec, float *itr, int *options, int *edgecut, idxtype *part,
                           MPI_Comm *comm)
```

### Description

This routine is used to balance the work load of a graph that corresponds to an adaptively refined mesh.

### Parameters

- vtxdist** This array describes how the vertices of the graph are distributed among the processors. (See discussion in Section 4.1). Its contents are identical for every processor.
- xadj, adjncy** These store the (local) adjacency structure of the graph at each processor. (See discussion in Section 4.1).
- vwgt, adjwgt** These store the weights of the vertices and edges. (See discussion in Section 4.1).
- vsize** This array stores the size of the vertices with respect to redistribution costs. Hence, vertices associated with mesh elements that require a lot of memory will have larger corresponding entries in this array. Otherwise, this array is similar to the vwgt array. (See discussion in Section 4.1).
- wgtflag** This is used to indicate if the graph is weighted. *wgtflag* can take one of four values:
- 0 No weights (vwgt and adjwgt are both NULL).
  - 1 Weights on the edges only (vwgt is NULL).
  - 2 Weights on the vertices only (adjwgt is NULL).
  - 3 Weights on both the vertices and edges.
- numflag** This is used to indicate the numbering scheme that is used for the *vtxdist*, *xadj*, *adjncy*, and *part* arrays. *numflag* can take the following two values:
- 0 C-style numbering is assumed that starts from 0
  - 1 Fortran-style numbering is assumed that starts from 1
- ncon** This is used to specify the number of weights that each vertex has. It is also the number of balance constraints that must be satisfied.
- nparts** This is used to specify the number of sub-domains that are desired. Note that the number of sub-domains is independent of the number of processors that call this routine.
- tpwgts** An array of size *ncon* x *nparts* that is used to specify the fraction of vertex weight that should be distributed to each sub-domain for each balance constraint. If all of the sub-domains are to be of the same size for every vertex weight, then each of the *ncon* x *nparts* elements should be set to a value of 1/*nparts*. If *ncon* is greater than one, the target sub-domain weights for each sub-domain are stored contiguously (similar to the vwgt array). Note that the sum of all of the *tpwgts* for a give vertex weight should be one.
- ubvec** An array of size *ncon* that is used to specify the imbalance tolerance for each vertex weight, with 1 being perfect balance and *nparts* being perfect imbalance. A value of 1.05 for each of the *ncon* weights is recommended.
- itr** This parameter describes the ratio of inter-processor communication time compared to data redistribution time. It should be set between 0.000001 and 1000000.0. If ITR is set high, a repartitioning with a low edge-cut will be computed. If it is set low, a repartitioning that requires little data redistribution will be computed. Good values for this parameter can be obtained by dividing inter-processor communication time by data redistribution time. Otherwise, a value of 1000.0 is recommended.

<b>options</b>	This is an array of integers that is used to pass additional parameters for the routine. If <i>options[0]</i> =0, then the default values are used. If <i>options[0]</i> =1, then the remaining three elements of <i>options</i> are interpreted as follows:
options[1]	This specifies the level of information to be returned during the execution of the algorithm. Timing information can be obtained by setting this to 1. Additional options for this parameter can be obtained by looking at the file <code>defs.h</code> in the <code>ParMETIS-Lib</code> directory. The numerical values there should be added to obtain the correct value. The default value is 0.
options[2]	This is the random number seed for the routine. The default value is 15.
options[3]	This specifies whether the sub-domains and processors are coupled or de-coupled. If the number of sub-domains desired ( <i>i.e.</i> , <code>nparts</code> ) and the number of processors that are being used is not the same, then these must be de-coupled. However, if <code>nparts</code> equals the number of processors, these can either be coupled or de-coupled. If sub-domains and processors are coupled, then the initial partitioning will be obtained implicitly from the graph distribution. However, if sub-domains are de-coupled from processors, then the initial partitioning needs to be obtained from the initial values assigned to the <code>part</code> array. A value of 1 indicates that sub-domains and processors are coupled and 2 indicates that these are de-coupled. The default value is 1 (coupled) if <code>nparts</code> equals the number of processors and 2 (de-coupled) otherwise.
<b>edgecut</b>	Upon successful completion, the number of edges that are cut by the partitioning is written to this parameter.
<b>part</b>	This is an array of size equal to the number of locally-stored vertices. Upon successful completion the partition vector of the locally-stored vertices is written to this array. (See discussion in Section 4.4). If the number of processors does not equal the number of sub-domains and/or options[3] is set to 2, then the previously computed partitioning must be passed to the routine as a parameter via this array.
<b>comm</b>	This is a pointer to the MPI communicator of the processes that call <code>PARMETIS</code> . For most programs this will point to <code>MPI_COMM_WORLD</code> .

### 5.3 Partitioning Refinement

**ParMETIS\_V3\_RefineKway** (idxtype \*vtxdist, idxtype \*xadj, idxtype \*adjncy, idxtype \*vwgt, idxtype \*adjwgt,  
int \*wgtflag, int \*numflag, int \*ncon, int \*nparts, float \*tpwgts, float \*ubvec,  
int \*options, int \*edgecut, idxtype \*part, MPI\_Comm \*comm)

#### Description

This routine is used to improve the quality of an existing  $k$ -way partitioning on  $p$  processors using the multi-level  $k$ -way refinement algorithm.

#### Parameters

- vtxdist** This array describes how the vertices of the graph are distributed among the processors. (See discussion in Section 4.1). Its contents are identical for every processor.
- xadj, adjncy** These store the (local) adjacency structure of the graph at each processor. (See discussion in Section 4.1).
- vwgt, adjwgt** These store the weights of the vertices and edges. (See discussion in Section 4.1).
- ncon** This is used to specify the number of weights that each vertex has. It is also the number of balance constraints that must be satisfied.
- nparts** This is used to specify the number of sub-domains that are desired. Note that the number of sub-domains is independent of the number of processors that call this routine.
- wgtflag** This is used to indicate if the graph is weighted. *wgtflag* can take one of four values:
- 0 No weights (vwgt and adjwgt are both NULL).
  - 1 Weights on the edges only (vwgt is NULL).
  - 2 Weights on the vertices only (adjwgt is NULL).
  - 3 Weights on both the vertices and edges.
- numflag** This is used to indicate the numbering scheme that is used for the *vtxdist*, *xadj*, *adjncy*, and *part* arrays. *numflag* can take the following two values:
- 0 C-style numbering is assumed that starts from 0
  - 1 Fortran-style numbering is assumed that starts from 1
- tpwgts** An array of size *ncon* x *nparts* that is used to specify the fraction of vertex weight that should be distributed to each sub-domain for each balance constraint. If all of the sub-domains are to be of the same size for every vertex weight, then each of the *ncon* x *nparts* elements should be set to a value of  $1/nparts$ . If *ncon* is greater than one, the target sub-domain weights for each sub-domain are stored contiguously (similar to the *vwgt* array). Note that the sum of all of the *tpwgts* for a give vertex weight should be one.
- ubvec** An array of size *ncon* that is used to specify the imbalance tolerance for each vertex weight, with 1 being perfect balance and *nparts* being perfect imbalance. A value of 1.05 for each of the *ncon* weights is recommended.
- options** This is an array of integers that is used to pass parameters to the routine. Their meanings are identical to those of **ParMETIS\_V3\_PartKway**.
- edgecut** Upon successful completion, the number of edges that are cut by the partitioning is written to this parameter.
- part** This is an array of size equal to the number of locally-stored vertices. Upon successful completion the partition vector of the locally-stored vertices is written to this array. (See discussion in Section 4.4).
- comm** This is a pointer to the MPI communicator of the processes that call **PARMETIS**. For most programs this will point to **MPI\_COMM\_WORLD**.

## 5.4 Fill-reducing Orderings

**ParMETIS\_V3\_NodeND** (idxtype \*vtxdist, idxtype \*xadj, idxtype \*adjncy, int \*numflag, int \*options,  
idxtype \*order, idxtype \*sizes, MPI\_Comm \*comm)

### Description

This routine is used to compute a fill-reducing ordering of a sparse matrix using multilevel nested dissection.

### Parameters

- vtxdist** This array describes how the vertices of the graph are distributed among the processors. (See discussion in Section 4.1). Its contents are identical for every processor.
- xadj, adjncy** These store the (local) adjacency structure of the graph at each processor (See discussion in Section 4.1).
- numflag** This is used to indicate the numbering scheme that is used for the *vtxdist*, *xadj*, *adjncy*, and *order* arrays. *numflag* can take the following two values:
- 0 C-style numbering is assumed that starts from 0
  - 1 Fortran-style numbering is assumed that starts from 1
- options** This is an array of integers that is used to pass parameters to the routine. Their meanings are identical to those of **ParMETIS\_V3.PartKway**.
- order** This array returns the result of the ordering (described in Section 4.4).
- sizes** This array returns the number of nodes for each sub-domain and each separator (described in Section 4.4).
- comm** This is a pointer to the MPI communicator of the processes that call **PARMETIS**. For most programs this will point to **MPI\_COMM\_WORLD**.

### Note

**ParMETIS\_V3.NodeND** requires that the number of processors be a power of 2.

## 5.5 Mesh to Graph Translation

**ParMETIS\_V3\_Mesh2Dual** (idxtype \*elmdist, idxtype \*eptr, idxtype \*eind, int \*numflag,  
int \*ncommonnodes, idxtype \*\*xadj, idxtype \*\*adjncy, MPI\_Comm \*comm)

### Description

This routine is used to construct a distributed graph given a distributed mesh. It can be used in conjunction with other routines in the **PARMETIS** library. The mesh can contain elements of different types.

### Parameters

**elmdist** This array describes how the elements of the mesh are distributed among the processors. It is analogous to the `vtxdist` array. Its contents are identical for every processor. (See discussion in Section 4.3).

**eptr, eind**

These arrays specifies the elements that are stored locally at each processor. (See discussion in Section 4.3).

**numflag** This is used to indicate the numbering scheme that is used for the *elmdist*, *elements*, *xadj*, *adjncy*, and *part* arrays. *numflag* can take one of two values:

0 C-style numbering that starts from 0.

1 Fortran-style numbering that starts from 1.

**ncommonnodes**

This parameter determines the degree of connectivity among the vertices in the dual graph. Specifically, an edge is placed between any two elements if and only if they share at least this many nodes. This value should be greater than zero, and for most meshes a value of two will create reasonable dual graphs. However, depending on the type of elements in the mesh, values greater than two may also be valid choices. For example, for meshes containing only triangular, tetrahedral, hexahedral, or rectangular elements, this parameter can be set to two, three, four, or two, respectively.

Note that setting this parameter to a small value will increase the number of edges in the resulting dual graph and the corresponding partitioning time.

**xadj, adjncy**

Upon the successful completion of the routine, pointers to the constructed *xadj* and *adjncy* arrays will be written to these parameters. (See discussion in Section 4.1).

**comm** This is a pointer to the MPI communicator of the processes that call **PARMETIS**. For most programs this will point to `MPI_COMM_WORLD`.

### Note

This routine can be used in conjunction with **ParMETIS\_V3\_PartKway**, **ParMETIS\_V3\_PartGeomKway**, or **ParMETIS\_V3\_AdaptiveRepart**. It typically runs in half the time required by **ParMETIS\_V3\_PartKway**.

## 6 Hardware & Software Requirements, and Contact Information

**PARMETIS** is written in ANSI C and uses MPI for inter-processor communication. Instructions on how to build **PARMETIS** are available in the `INSTALL` file. In the directory called `Graphs`, you will find programs that tests if **PARMETIS** was built correctly. Also, a header file called `parmetis.h` is provided that contains prototypes for the functions in **PARMETIS**.

In order to use **PARMETIS** in your application you need to have a copy of the serial **METIS** library and link your program with both libraries (*i.e.*, `libparmetis.a` and `libmetis.a`). Note that the **PARMETIS** package already contains the source code for the **METIS** library. The included Makefiles automatically construct both libraries.

**PARMETIS** have been extensively tested on a number of different parallel computers. However, even though **PARMETIS** contains no known bugs, this does not mean that all of its bugs have been found and fixed. If you have any problems, please send email to `metis@cs.umn.edu` with a brief description of the problem.

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